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APPLICATION NO.	FILING DATE	FIRST NAMED INVENTOR	ATTORNEY DOCKET NO.	CONFIRMATION NO.	
10/722,374	11/25/2003	David Bebbington	VPI/00-130-08 CON US	VPI/00-130-08 CON US 8573	
27916	7590 04/11/2005		EXAMINER		
VERTEX PHARMACEUTICALS INC.			BALASUBRAMANIAN	BALASUBRAMANIAN, VENKATARAMAN	
130 WAVERLY STREET CAMBRIDGE, MA 02139-4242			ART UNIT	PAPER NUMBER	
			1624		
			DATE MAILED: 04/11/200:	5	

Please find below and/or attached an Office communication concerning this application or proceeding.

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	·	Application No.	Applicant(s)	_		
		10/722,374	BEBBINGTON ET AL.			
•	Office Action Summary	Examiner	Art Unit			
		Venkataraman Balasubramanian	1624			
Period fo	<ul> <li>The MAILING DATE of this communication app or Reply</li> </ul>	pears on the cover sheet with the c	orrespondence address			
THE - Exte after - If th - If NC - Failt Any	MAILING DATE OF THIS COMMUNICATION.  Insions of time may be available under the provisions of 37 CFR 1.13  IT SIX (6) MONTHS from the mailing date of this communication.  It period for reply specified above is less than thirty (30) days, a reply of period for reply is specified above, the maximum statutory period we ure to reply within the set or extended period for reply will, by statute, reply received by the Office later than three months after the mailing and patent term adjustment. See 37 CFR 1.704(b).	36(a). In no event, however, may a reply be time within the statutory minimum of thirty (30) days will apply and will expire SIX (6) MONTHS from a cause the application to become ABANDONE	nely filed s will be considered timely. the mailing date of this communication. D (35 U.S.C. § 133).			
Status						
1) 又	Responsive to communication(s) filed on 18 Ja	anuary 2005.				
		action is non-final.				
3)	Since this application is in condition for allowance except for formal matters, prosecution as to the merits is closed in accordance with the practice under <i>Ex parte Quayle</i> , 1935 C.D. 11, 453 O.G. 213.					
Disposit	ion of Claims	,				
5)□ 6)⊠ 7)□	Claim(s) <u>1-26</u> is/are pending in the application.  4a) Of the above claim(s) is/are withdraw Claim(s) is/are allowed.  Claim(s) <u>1-26</u> is/are rejected.  Claim(s) is/are objected to.  Claim(s) are subject to restriction and/or	vn from consideration.				
Applicat	ion Papers					
9)[	The specification is objected to by the Examine	r.	•			
10)	The drawing(s) filed on is/are: a) acc	epted or b)□ objected to by the I	Examiner.			
	Applicant may not request that any objection to the	drawing(s) be held in abeyance. See	e 37 CFR 1.85(a).			
11)	Replacement drawing sheet(s) including the correct The oath or declaration is objected to by the Ex	• • • • • • • • • • • • • • • • • • • •				
Priority (	under 35 U.S.C. § 119		•			
12)□ a)	Acknowledgment is made of a claim for foreign  All b) Some * c) None of:  1. Certified copies of the priority documents  2. Certified copies of the priority documents  3. Copies of the certified copies of the priority application from the International Bureau  See the attached detailed Office action for a list	s have been received. s have been received in Applicati nty documents have been receive u (PCT Rule 17.2(a)).	on No ed in this National Stage			
Attachmen	ut(s)		•			
1) Notice	ce of References Cited (PTO-892)	4) Interview Summary	(PTO-413)			
3) 🛛 Infor	ce of Draftsperson's Patent Drawing Review (PTO-948) mation Disclosure Statement(s) (PTO-1449 or PTO/SB/08) er No(s)/Mail Date 3/22/04.	Paper No(s)/Mail Da		Ì		

U.S. Patent and Trademark Office PTOL-326 (Rev. 1-04)

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### **DETAILED ACTION**

### Election/Restrictions

Applicant's election of Group I, claims 1-26 drawn to compound of formula III wherein  $Z^1$  or  $Z^3$  is nitrogen the other  $Z^1$  or  $Z^3$  is  $CR^8$  or  $CR^x$  respectively and  $Z^2$  is CH, in the reply filed on 1/18/2005 is acknowledged. Because applicant did not distinctly and specifically point out the supposed errors in the restriction requirement, the election has been treated as an election without traverse (MPEP § 818.03(a)).

Claims 1-26 are pending.

## Information Disclosure Statement

References cited in the Information Disclosure Statement, filed on 3/22/2004, are made of record.

### Specification

Specification needs to be amended to indicate this application is a continuation of US Application No. 10/034,683, filed 12/20/2001 now US Patent No. 6,656,939.

### Claim Rejections - 35 USC § 112

The following is a quotation of the second paragraph of 35 U.S.C. 112:

The specification shall conclude with one or more claims particularly pointing out and distinctly claiming the subject matter which the applicant regards as his invention.

Claims 1-26 are rejected under 35 U.S.C. 112, second paragraph, as being indefinite for failing to particularly point out and distinctly claim the subject matter which applicant regards as the invention. Following reasons apply. Any claim not specifically rejected is rejected as being dependent on a rejected claim.

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1. Recitation of "pharmaceutically acceptable derivative or prodrug thereof" in claim 1 and 2 renders these claims indefinite as it implies more than what is being positively recited. The term derivative can include any organic compound and hence it is not clear what is being claimed. The metes and bounds of the these derivative therefore remains unknown. In addition, claims 1 and 2 also include prodrug. Prodrugs in general and as noted in specification, are compounds, which undergo in vivo hydrolysis to parent active drugs. In that sense recitation of prodrugs is acceptable. However, the definition of pharmaceutical derivatives (page 23) appears to include various functional groups include such groups, namely esters, etc. and therefore it is not clear what is the difference between these variable groups and the prodrug groups. In addition, reading the specification (page 24), the phrase is meant to include without limitation esters, amino acid esters, phosphate esters metal salts and sulfonate esters. Again the definition of various varible group embraced for compound of formula I include such esters and therefore it is not clear how would distinguish esters as variable groups and esters as prodrugs. Furthermore, the issue on second paragraph is whether the structures of the claimed compounds are clearly defined. Applicants' "prodrugs" are molecules whose structure lie outside the subject matter of formula (I), but upon metabolism in the body are converted to active compounds falling within the structural scope of formula (I). The claim describes the function intended but provides no specific structural guidance to what constitutes a

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"prodrug". Structural formulas, names, or both can accurately describe organic compounds, which are the subject matter of claim 1. Attempting to define means by function is not proper when the means can be clearly expressed in terms that are more precise. Applicants list the function that these prodrugs are to perform in the passage spanning line 11-14 of page 24 but offer no structural guidance as to which derivatives are intended.

2. Recitation of "one or more features" in claims 5 and 7, renders these claims indefinite, as it is not clear what is considered as "features" in the recited groups therein to be selected.

The following is a quotation of the first paragraph of 35 U.S.C. 112:

The specification shall contain a written description of the invention, and of the manner and process of making and using it, in such full, clear, concise, and exact terms as to enable any person skilled in the art to which it pertains, or with which it is most nearly connected, to make and use the same and shall set forth the best mode contemplated by the inventor of carrying out his invention.

Claims 1-26 are rejected under 35 U.S.C. 112, first paragraph, because the specification, while being enabling for making salts of the claimed compounds, does not reasonably provide enablement for making prodrugs of the claimed compounds. The claim(s) contains subject matter that was not described in the specification in such a way as to enable one skilled in the art of medicinal chemistry - to use the invention. "The factors to be considered in making an enablement rejection have been summarized as the quantity of experimentation necessary, the amount of direction or guidance presented, the presence or absence of working examples, the nature of the invention, the state of the prior art, the relative skill of those in that art, the predictability or unpredictability of the

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art and the breadth of the claims", In re Rainer, 146 USPQ 218 (1965); In re Colianni, 195 USPQ 150, Ex parte Formal, 230 USPQ 546. a) Finding a prodrug is an empirical exercise. Predicting if a certain ester of a claimed alcohol, for example, is in fact a prodrug, and produces the active compound metabolically, in man, at a therapeutic concentration and at a useful rate is filled with experimental uncertainty. Although attempts have been made to predict drug metabolism 'de novo, this is still an experimental science. For a compound to be a prodrug, it must meet three tests. It must itself be biologically inactive. It must be metabolized to a second substance in a human at a rate and to an extent to produce that second substance at a physiologically meaningful concentration. Thirdly, that second substance must be biologically active. Thus, determining whether a particular compound meets these three criteria in a clinical trial setting requires a large quantity of experimentation.

b) The direction concerning the prodrugs is found in the passage spanning line 11-14 of page 24 c) There is no working example of a prodrug of a compound the formula (I). d) The nature of the invention is clinical use of compounds and the pharmacokinetic behavior of substances in the human body. e) The state of the prodrug art is summarized by Wolff (Medicinal Chemistry). The table on the left side of page 976 outlines the research program to be undertaken to. find a prodrug. The second paragraph in section 10 and the paragraph spanning pages 976-977 indicate the low expectation of success. In that paragraph the difficulties of extrapolating between species are further developed. Since, the prodrug concept is a pharmacokinetic issue, the lack of any standard pharmacokinetic

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protocol discussed in the last sentence of this paragraph is particularly relevant. Banker (Modem Pharmaceutics) in the first sentence, third paragraph on page 596 states that "extensive development must be undertaken" to find a prodrug. I) Wolff (Medicinal Chemistry) in the last paragraph on page 975 describes the artisans making Applicants' prodrugs as a collaborative team of synthetic pharmaceutical chemists and metabolism experts. All would have a Ph. D. degree and several years of industrial experience. g) It is well established that "the scope of enablement varies inversely degree of unpredictability of the factors involved", 'and physiological activity is generally considered to be an unpredictable factor. See In re Fisher, 427 F.2d 833, 839, 166 USPQ 18, 24 (CCPA 1970). h) The breadth of the claims includes all of the hundreds of thousands of compounds of formula of claim I as well as the presently unknown list potential prodrug derivatives embraced by the word "prodrug".

Thus, undue experimentation will be required to determine if any particular derivative is, in fact, a prodrug.

MPEP §2164.01(a) states, "A conclusion of lack of enablement means that, based on the evidence regarding each of the above factors, the specification, at the time the application was 'filed, would not have taught one skilled in the art how to make and/or use the full scope of the claimed invention without undue experimentation. In re Wright, 999 F.2d 1557,1562, 27 USPQ2d 1510, 1513 (Fed. Cir. 1993)." That conclusion is clearly justified here and undue experimentation will be required to practice Applicants' invention.

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Claims 13-22 and 24-26 are rejected under 35 U.S.C. 112, first paragraph, because the specification, while being enabling for treating diabetes does not reasonably provide enablement for treatment any or all diseases including those yet to be linked with the various mode of action embraced in the claim language. The specification does not enable any person skilled in the art to which it pertains, or with which it is most nearly connected, to use the invention commensurate in scope with these claims.

The instant claims are drawn to a method of inhibiting or treating a disease in a patient of based on the mode of action as Aurora-2, GSK-3 and phosphoryrlation of protein (Tau or β-catenin which as recited reads on any or all diseases of these organs for which there is no enabling disclosure. These claims are deemed as reach through claims wherein based on the mode of action, treating any or all diseases is embraced. The scope of the claims includes treatment of any or all of diseases, which are not adequately enabled solely based on the activity of the compounds, provided in the specification at pages 69-70. The instant compounds are disclosed have Aurora-2, GSK-3 inhibitory activity & inhibition of phosphoryrlation of protein. It is therefore recited that the instant compounds are useful in treating any or all diseases for which applicants provide no competent evidence. Reading specification it appears that instant compound is useful for treating all sorts of diseases for which applicants have not provided any experimental support or nexus. Prior art search and those cited in the Information disclosure statement do not lend support to, except for treating diabetes, treatments of all diseases embraced in the claim language. That a

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single class of compounds can treat all or any disease of the said organs is an incredible finding for which applicants have not provided enabling disclosure and the Information Disclosure Statement suggest the use of these inhibitors is still under experimental stage and speculative in nature. Prior art search also shows that further studies are needed. See Kim et al. Differentiation and Gene Regulation 508-514, 2000 and Warner et al., Molecular Cancer Therapeutics 2: 589-595, 2003. Furthermore, the applicants have not provided any competent evidence that the instantly disclosed tests are highly predictive for all the uses disclosed and embraced by the claim language for the intended host. Moreover many if not most of diseases such as Alzheimer's disease, multiple sclerosis, ALS etc. are very difficult to treat and at present there is no known drug, which can successfully reverse the course of these diseases, despite the fact that there are many drugs, which can be used for "inflammatory condition". The scope of the claims involves all of the thousands of compounds of claim 1 as well as the thousand of diseases embraced by the terms Aurora-2 mediated disease and or GSK-3mediated disease.

No compound has ever been found to treat cancers of all types generally. Since this assertion is contrary to what is known in medicine, proof must be provided that this revolutionary assertion has merits. The existence of such a "compound" is contrary to our present understanding of pathology of disease and their treatment.

Note substantiation of utility and its scope is required when utility is "speculative", "sufficiently unusual" or not provided. See Ex parte Jovanovics,

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211 USPQ 907, 909; In re Langer 183 USPQ 288. Also note Hoffman v. Klaus 9 USPQ 2d 1657 and Ex parte Powers 220 USPQ 925 regarding type of testing needed to support in vivo uses.

Next, applicant's attention is drawn to the Revised Utility and Written Description Guidelines, at 66 FR 1092-1099, 2001 wherein it is emphasized that 'a claimed invention must have a specific and substantial utility'. The disclosure in the instant case is not sufficient to enable the instantly claimed method treating solely based on the inhibitory activity disclosed for the compounds. The state of the art is indicative of the requirement for undue experimentation. See Kim et al. Differentiation and Gene Regulation 508-514, 2000 and Warner et al., Molecular Cancer Therapeutics 2: 589-595, 2003.

In evaluating the enablement question, several factors are to be considered. Note *In re Wands*, 8 USPQ2d 1400 and *Ex parte Forman*, 230 USPQ 546. The factors include: 1) The nature of the invention, 2) the state of the prior art, 3) the predictability or lack thereof in the art, 4) the amount of direction or guidance present, 5) the presence or absence of working examples, 6) the breadth of the claims, and 7) the quantity of experimentation needed.

- 1) The nature of the invention: Therapeutic use of the compounds in treating several diseases of that require Aurora-2 a and GSK-3 inhibitory activity as well as inhibition of phosphoryrlation of protein
- 2) The state of the prior art: A recent publication in the Information Disclosure Statement as well as in the specification suggest that Aurora-2, and GSK-3 are still in experimental stage. See Kim et al. Differentiation and Gene Regulation

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508-514, 2000 and Warner et al., Molecular Cancer Therapeutics 2: 589-595, 2003.

- 3) The predictability or lack thereof in the art: Applicants have not provided any competent evidence or disclosed tests that are highly predictive for the pharmaceutical use for the therapeutic effect of the instant compounds. Pharmacological activity in general is a very unpredictable area. Note that in cases involving physiological activity such as the instant case, "the scope of enablement obviously varies inversely with the degree of unpredictability of the factors involved". See *In re Fisher*, 427 F.2d 833, 839, 166 USPQ 18, 24 (CCPA 1970).
- 4) The amount of direction or guidance present and 5) the presence or absence of working examples: Specification has no working examples to show therapeutic effect and the state of the art is that the effects of Aurora-2, GSK-3 inhibitors and phosphorylation of Tau protein inhibitors are still in experimental stage
- 6) The breadth of the claims: The instant claims embrace treatment of several diseases including those yet to be related to Aurora-2 and GSK-3 activity.
- 7) The quantity of experimentation needed would be an undue burden to one skilled in the pharmaceutical arts since there is inadequate guidance given to the skilled artisan, regarding the pharmaceutical use, for the reasons stated above. Thus, factors such as "sufficient working examples", "the level of skill in the art" and "predictability", etc. have been demonstrated to be sufficiently lacking in the instant case for the instant method claims. In view of the breadth of the claims, the chemical nature of the invention, the unpredictability of ligand-receptor

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interactions in general, and the lack of working examples regarding the activity of the claimed compounds towards treating the variety of diseases of the instant claims, one having ordinary skill in the art would have to undergo an undue amount of experimentation to use the instantly claimed invention commensurate in scope with the claims.

MPEP §2164.01(a) states, "A conclusion of lack of enablement means that, based on the evidence regarding each of the above factors, the specification, at the time the application was 'filed, would not have taught one skilled in the art how to make and/or use the full scope of the claimed invention without undue experimentation. In re Wright, 999 F.2d 1557,1562, 27 USPQ2d 1510, 1513 (Fed. Cir. 1993)." That conclusion is clearly justified here and undue experimentation will be required to practice Applicants' invention.

#### Conclusion

Any inquiry concerning this communication from the examiner should be addressed to Venkataraman Balasubramanian (Bala) whose telephone number is (571) 272-0662. The examiner can normally be reached on Monday through Thursday from 8.00 AM to 6.00 PM. The Supervisory Patent Examiner (SPE) of the art unit 1624 is Mukund Shah whose telephone number is (571) 272-0674. If Applicants are unable to reach Mukund Shah within 24-hour period, they may contact James O. Wilson, Acting-SPE of art unit 1624 at 571-272-0661.

The fax phone number for the organization where this application or proceeding is assigned (703) 872-9306. Any inquiry of a general nature or

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relating to the status of this application or proceeding should be directed to the receptionist whose telephone number is (571) 272-1600.

Information regarding the status of an application may be obtained from the Patent Application Information Retrieval (PAIR) system. Status information for published applications may be obtained from either Private PAIR or Public PAG. Status information for unpublished applications is available through Private PAIR only. For more information about the PAIR system, see http://pair-direct.uspto.gov. Should you have questions on access to the Private PAIR system, contact the Electronic Business Center (EBC) at 866-2 17-9197 (toll-free).

Venkataranan Balasubramuhan Venkataraman Balasubramanian

4/1/2005